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## Simplified Free-Energy Calculation for Ion-Induced Heteromolecular Nucleation

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# Simplified Free-Energy Calculation for Ion-Induced Heteromolecular Nucleation

Jag J. Singh and Alphonso C. Smith Langley Research Center Hampton, Virginia

Lo Y. Chan and Glenn K. Yue Institute for Atmospheric Optics and Remote Sensing Hampton, Virginia



Scientific and Technical Information Branch

#### SUMMARY

A modified expression for the Gibbs free energy of formation of ion-induced microclusters in binary vapor mixtures has previously been developed by including the effects of microscopic surface tension, field-dependent, nonlinear, dielectric behavior of liquids, and neutral clustering prior to ion-induced nucleation. It has been further simplified by using an empirical correction factor for the electrostatic term instead of the tedious, time-consuming computations required previously. The electrostatic correction factor has been obtained by comparing the simplified-theory calculations for free energy of selected ion-hydrate formation with more rigorous molecular-dynamics calculations. The simplified Gibbs free-energy expression has been applied to calculate the spectral distribution of ion-H2O-H2SO4 microclusters for various combinations of relative humidity and relative acidity. Because of the higher contribution of the electrostatic energy term in the simplified expression, the predicted microcluster spectra are broader and contain more of the larger ion clusters than predicted by the earlier theories. These trends are consistent with recently reported experimental data on ion-induced binary nucleation.

#### INTRODUCTION

It is well-known that ions are present throughout the atmosphere. In the upper atmosphere, galactic cosmic rays, solar flares, and ultraviolet radiation from the Sun induce ionization of gaseous molecules. Closer to the surface of the Earth, radioactivity associated with uranium/thorium series ores as well as coal combustion products help to produce primary ions. The ions act as foreign centers and greatly enhance the nucleation process. They facilitate the formation of stable, ultrafine, prenucleation clusters, especially in the presence of traces of pollutant gases. Rocket measurements show that stable ion clusters are part of the aerosol size spectrum in the stratosphere. (See refs. 1 and 2.) Ion-induced gas-to-particle conversion plays an important role in the formation of ultrafine aerosols emitted from high-temperature combustion sources. (See ref. 3.) Thus, a study of the mechanism of ion-induced nucleation is very important in gaining a better understanding of space and Earth environments. In the present study, we address the problem of nucleation processes induced by the presence of ions in a binary mixture of vapors. An empirical modification of the classical nucleation theory is presented, and the cluster spectra obtained using this modified approach are compared with other existing theories.

The conventional Thomson theory (ref. 4) for hydration of ions is developed from the ion-liquid drop theory and hence suffers from all the shortcomings of the liquid drop model. (See refs. 5 through 10.) According to this theory, the free energy of formation of ion clusters is composed of three parts: (1) Volume energy, (2) surface energy, and (3) electrostatic energy. There are two major criticisms of the classical ion-liquid droplet theory. First, bulk values are used in the volume and surface-energy terms, even for small clusters. Second, the electrostatic-energy term is treated by considering the ion as embedded in a uniform dielectric medium and thus neglecting the strong intermolecular interaction between the ion core and its surrounding molecules. To improve the conventional theory, Chan and Mohnen (ref. 9) developed a semimolecular ion-nucleation theory which includes corrections for the aforementioned defects. Briant and Burton (ref. 11) adopted a molecular approach

that is partially based on molecular dynamics. Some of the corrections proposed in reference 9 have been extended further so that the theory can apply to ion-induced nucleation in binary mixtures of vapors. (See ref. 12.)

It is the purpose of this paper to show that use of the microscopic surface tension and appropriate scaling of the electrostatic-energy term in the Thomson theory for ion hydrates can produce good agreement with results obtained with the more rigorous Chan and Mohnen (ref. 9) and Briant and Burton (ref. 11) models. The simple scaling corrections thus obtained are then extended to the ion-induced nucleation in binary mixtures of vapors. Predicted results for the formation of ion-H<sub>2</sub>O-H<sub>2</sub>SO<sub>4</sub> clusters based on the Thomson theory for binary mixtures of vapors (model I), the more complicated modified theory (ref. 12) for ion-induced binary nucleation (model II), and the present modified Thomson theory (model III), are compared, and the implications are discussed in the following sections.

#### SYMBOLS

Е	electric field, $\frac{Q}{\epsilon(r)r^2}$
ΔG	free energy of droplet formation
ΔGB	bulk energy term
ΔG <sub>E</sub>	electrostatic energy term
ΔG	free-energy change without effect of formation of sulfuric acid hydrate
ΔGs	surface energy term
g	field-dependent Kirkwood factor
k	Boltzmann constant
N <sub>A,B</sub>	total number of molecules of component A or B in system
n <sub>A,B</sub>	number of molecules of component A (water) or B (sulfuric acid) in cluster
P(n <sub>A</sub> ,n <sub>B</sub> )	probability of occurrence for the ion cluster containing $n_{\! A}^{}$ molecules of component B $$
PA,B	equilibrium vapor pressure of component A or B
$p_{A}^{O}$	saturation vapor pressure of component A
$p_{B}^{O}$	saturation vapor pressure of component B
$p_{A,B}^{sol}$	equilibrium vapor pressure of component A or B over flat surface of solution
p,	partial pressure of component A or B after sulfuric acid hydrate formation
Q	ionic charge on cluster

 $Q_{\mbox{\scriptsize eff}}$  effective ionic charge on cluster

r radius of droplet

r<sub>1</sub> ionic radius

S saturation ratio

S<sub>A</sub> relative humidity (R.H. in tables)

 $S_R$  relative acidity (R.A. in tables)

T absolute temperature

 $\alpha$  empirically adjustable scaling parameter

 $\alpha_n$  water activity

 $\alpha_{_{\rm B}}$  acid activity

$$\beta = \frac{3}{2kT} \left( \frac{\eta^2 + 2}{3} \right) \mu_0 g^{1/2}$$

γ charge scaling factor

δ Tolman coefficient

ε dielectric constant of mixture

 $\epsilon_{\gamma}$  bulk dielectric constant of mixture

ε(r) radius-dependent dielectric constant

η internal refractive index of liquid

 $\mu_{\Omega}$  dipole moment in absence of field

σ surface tension

o plane surface tension

#### THEORETICAL CONSIDERATIONS

The classical Thomson theory states that the free energy of formation  $\Delta G$  for an ion-hydrate cluster is given by

$$\Delta G = \Delta G_{B} + \Delta G_{S} + \Delta G_{E}$$
 (1)

The bulk-energy term is given by

$$\Delta G_{R} = -n_{A}kT \ln S \tag{2}$$

the surface-energy term is

$$\Delta G_{S} = 4\pi r^{2} \sigma \tag{3}$$

and the electrostatic term is

$$\Delta G_{E} = \frac{Q^{2}}{2} \left( 1 - \frac{1}{\varepsilon} \right) \left( \frac{1}{r} - \frac{1}{r_{1}} \right) \tag{4}$$

where

number of molecules of component A

k Boltzmann constant

T absolute temperature

S saturation ratio (ratio of partial pressure to saturation pressure)

r radius of ion cluster

σ surface tension

Q ionic charge

ε dielectric constant

r<sub>1</sub> ionic radius

In the calculation of the surface-energy term (eq. (3)), the bulk surface-tension value is used, and in equation (4), the ion is assumed to be embedded in a uniform dielectric medium. Results obtained using the Thomson theory do not agree with the semimolecular theory developed by Chan and Mohnen (ref. 9) as well as with the molecular theory, based partially on molecular dynamics, developed by Briant and Burton (ref. 11). A major source of error is the use of the macroscopic surface tension and the neglect of the strong intermolecular interactions between the ion core and the surrounding water molecules. It is well recognized that the surface tension is curvature-dependent. (See ref. 13.) The influence of this curvature effect on the free energy of formation of microclusters has been discussed in several recent papers. (See refs. 13 and 14, for example.) The Tolman correction for the surface tension (ref. 13) is simple and widely used. Using the Tolman coefficient  $\delta$ , we have

$$\sigma(r) = \sigma_0 \left( 1 + \frac{2\delta}{r} \right)^{-1} \tag{5}$$

where  $\sigma(r)$  is the surface tension of a microcluster of radius r and  $\sigma$  is the plane surface tension. Vogelsberger (ref. 15) suggested 0.92 Å as the value of  $\delta$  for water.

The correction for the electrostatic term is more complicated. It takes into account the influence of the intermolecular interaction between the ion core and the surrounding molecules as well as the breakdown of the linear correlation between dielectric polarization and electric-field strength. Corrections for this defect are discussed by Chan and Mohnen (ref. 9). Briant and Burton (ref. 11) treat this problem by applying molecular-dynamics calculations. Since the computations for the free energy of formation of gas-phase ion hydrates by Chan and Mohnen (ref. 9) and Briant and Burton (ref. 11) are among the most accurate, one simple approach to correct for this defect is to empirically scale the electrostatic term so that good agreement is reached between the present calculations and those of references 9 and 11. Incorporating the Tolman correction for the surface tension and the empirical scaling factor for the electrostatic term, the modified expression for the Gibbs free energy is

$$\Delta G = -n_A kT \ln S + 4\pi r^2 \sigma_O \left(1 + \frac{2\delta}{r}\right)^{-1} + \gamma \left(\frac{Q^2}{2}\right) \left(1 - \frac{1}{\epsilon}\right) \left(\frac{1}{r} - \frac{1}{r_1}\right)$$
 (6)

where  $\gamma$  is the charge scaling factor. The value of  $\Delta G$  is critically dependent on the value of  $\gamma_{\bullet}$ 

Free energies of F and Cs<sup>+</sup> hydrates have been calculated using equation (6). The initial hydrate growth is largely determined by the electrostatic interaction between the ion and the surrounding molecules and is, therefore, strongly dependent on the value of  $\gamma$ . A small change ( $\approx 10$  percent) in the value of  $\gamma$  can produce a measurable change in the ion-hydrate spectrum. Figures 1 and 2 show the comparison between various theories for the calculation of  $\Delta G$  for two ion hydrates. The agreement is especially good between the calculations of Briant and Burton and those of the present model for  $\gamma = 1.3$ . The scaling procedure suggests that when microscopic surface tension is taken into account, the effect of the interaction between the ion core and the surrounding molecules can be approximated by using an effective charge distribution of

$$Q_{eff} = \sqrt{1.3Q} = 1.14Q$$
 (7)

A value of  $Q_{\mbox{eff}}$  greater than Q implies relatively greater importance of the electrostatic term than of the free-energy change of the neutral portion of the cluster in the prenucleation stages.

Suck (ref. 16) has also calculated the contribution of the electrostatic interaction energy between the ion and the surrounding molecules in the cluster. In the

limiting case of zero polarizability and zero dipole moment associated with an ion, his expression for the electrostatic term reduces to the following form:

$$\Delta G_{\text{electrostatic}} = \frac{9}{2} \left[ \frac{\epsilon(\epsilon - 1)}{(2\epsilon + 1)(\epsilon + 2)} \right] \alpha Q^2 \left( \frac{1}{r} - \frac{1}{r_1} \right)$$

$$\approx \frac{9\alpha}{2} \left(\frac{Q^2}{2}\right) \left(1 - \frac{1}{\varepsilon}\right) \left(\frac{1}{r} - \frac{1}{r_1}\right) \tag{8a}$$

where  $\alpha$  is an empirically adjustable scaling parameter. Using equation (8a), Suck calculated free energy of formation of  $F^{-} \cdot (H_2O)_n$  and  $Cs^{+} \cdot (H_2O)_n$  clusters for direct comparison with computations of Briant and Burton (ref. 11). He found good agreement for  $\alpha = 1/3$ , thus giving

$$Q_{eff} = \sqrt{1.5Q} = 1.22Q$$
 (8b)

However, Suck's primary interest was to assess the importance of the electrostatic term to the total free-energy change owing to the presence of ions during the cluster formation. This was in contrast to the conventional Thomson theory, in which the electrostatic term is usually added to the free-energy change of a neutral drop to obtain the total free-energy change. Consequently, Suck had made no changes in the volume and the surface-energy terms (i.e., he did not include size dependence of the surface tension in his calculations).

Figures 1 and 2 show that the same value of the scaling factor (i.e.,  $\gamma$  = 1.3) applies equally well for the positive and negative ions and works fine for clusters containing up to 30 water molecules. The concept of the scaling factor for the electrostatic-energy term is further extended to the ion-induced nucleation in a binary mixture of vapors in the following section.

### COMPARISON OF DIFFERENT BINARY ION-NUCLEATION THEORIES FOR MIXTURE

#### OF WATER AND SULFURIC ACID VAPORS

When the Thomson theory is extended to binary ion nucleation for a mixture of vapors (ref. 17), it states that the free energy required to form a droplet  $\Delta G$  with  $n_A$  molecules of substance A and  $n_B$  molecules of substance B around an ion core with charge Q and radius  $r_1$  (model I) is given by the following equation:

$$\Delta G(n_{A}, n_{B}) = -n_{A}kT \ln \frac{S_{A}}{\alpha_{A}} - n_{B}kT \ln \frac{S_{B}}{\alpha_{B}} + 4\pi r^{2}\sigma_{O} + \frac{Q^{2}}{2}\left(1 - \frac{1}{\varepsilon_{O}}\right)\left(\frac{1}{r} - \frac{1}{r_{1}}\right)$$
(9)

where

$$S_{A} = \frac{P_{A}}{P_{A}} \tag{10a}$$

$$\alpha_{A} = \frac{p_{A}^{\text{sol}}}{p_{A}^{\text{o}}} \tag{10b}$$

$$S_{B} = \frac{P_{B}}{P_{B}}$$
 (10c)

$$\alpha_{\rm B} = \frac{p_{\rm B}^{\rm sol}}{p_{\rm B}^{\rm o}} \tag{10d}$$

where

p<sub>i</sub> equilibrium vapor pressure of specie i

p, saturation vapor pressure of pure specie i

 $_{
m i}^{
m sol}$  equilibrium vapor pressure of specie i over flat surface of solution

In this paper,  $n_{\rm A}$  and  $n_{\rm B}$  are used to denote the number of water and sulfuric acid molecules in the ion cluster, respectively.

A more complex equation for calculation of the free energy of droplet formation in binary mixtures has been developed by Singh et al. (See ref. 9.) The free energy of formation of the ion clusters (model II) takes the form

$$\Delta G(n_{A}, n_{B}) = -n_{A}kT \ln \frac{S_{A}}{\alpha_{A}} - n_{B}kT \ln \frac{S_{B}}{\alpha_{B}} + 4\pi r^{2}\sigma_{O}\left(1 + \frac{2\delta}{r}\right)^{-1}$$

$$- n_{B}kT \ln \frac{p_{B}'}{p_{B}} + \frac{Q^{2}}{2} \left(\frac{1}{r} - \frac{1}{r_{1}}\right) + \frac{Q^{2}}{2} \int_{r_{1}}^{r} \frac{1}{\epsilon(r)r^{2}} dr$$
 (11)

where  $\delta$  is the Tolman coefficient,  $p_B^{\prime}$  is the actual vapor pressure of the sulfuric acid taking into account the presence of the sulfuric acid hydrates, and  $\epsilon(r)$ 

is the radius-dependent dielectric constant. The first two terms on the right-hand side of equation (11) are the bulk energy terms. The third term represents the surface energy, taking into account the effect of curvature of small droplets. The fourth term accounts for the effect of the formation of neutral sulfuric acid hydrates. The value of  $p_B^*/p_B$  in this term is obtained by solving the following equation numerically (refs. 12 and 18):

$$\sum_{n_{A}} \sum_{n_{B}} N_{A} \left(\frac{p_{B}^{\prime}}{p_{B}}\right)^{n_{B}} \exp\left[-\frac{\Delta G_{O}(n_{A}, n_{B})}{kT}\right] = N_{B} \left(1 - \frac{p_{B}^{\prime}}{p_{B}}\right)$$
(12)

where  $N_A$  and  $N_B$  are the total number of molecules for A and B in the system and  $\Delta G_O$  is the free-energy change without the effect of the formation of sulfuric acid hydrate. The fifth and sixth terms in equation (11) describe the electrostatic-energy contribution. The value of  $\epsilon(r)$  can be obtained by solving the following equations (refs. 12 and 19):

$$\varepsilon(\mathbf{r}) = \eta^2 + \left(\varepsilon_0 - \eta^2\right) \frac{3}{\beta E} \left(\frac{1}{\tanh(\beta E)} - \frac{1}{\beta E}\right) \tag{13a}$$

where

$$E = \frac{Q}{\varepsilon(r)r^2} \tag{13b}$$

$$\beta = \frac{3}{2kT} \left( \frac{\eta^2 + 2}{3} \right) \mu_0 g^{1/2}$$
 (13c)

Here,  $\eta$  is the internal refractive index of the liquid,  $\mu_O$  is the dipole moment in the absence of the field, and g is the field-dependent Kirkwood factor (ref. 20) defined as follows:

$$g = g_0 + g_1 E^2 + g_2 E^4 + g_3 E^6 + \dots$$
 (13d)

If higher terms in E are neglected, then

$$g \approx 1 + \frac{g_0 - 1}{1 - \left[g_1 E^2 / (g_0 - 1)\right]}$$
 (13e)

The values of  $g_0$  and  $g_1$  for water given by Liszi et al. (ref. 19) are as follows:  $g_0 = 2.8$  and  $g_1 = -0.56 \times 10^{-8}$  (when E is expressed in esu).

Using a simplified approach to calculate the electrostatic-energy term, equation (11) can be modified as follows (model III):

$$\Delta G = -n_A kT \ln \frac{S_A}{\alpha_A} - n_B kT \ln \frac{S_B}{\alpha_B} + 4\pi r^2 \sigma_O \left(1 + \frac{2\delta}{r}\right)^{-1}$$

$$+ \gamma \left(\frac{Q^2}{2}\right) \left(1 - \frac{1}{\epsilon}\right) \left(\frac{1}{r} - \frac{1}{r_1}\right) - n_B kT \ln \frac{p_B^1}{p_B}$$
(14)

where  $\gamma = 1.3.**$ 

The value of surface tension  $\sigma$  and the variation of  $p_A^{sol}$  and  $p_B^{sol}$  with composition are available in the literature. (See refs. 21 and 22.)

A major advantage of model III over model II is that by empirically adjusting the electrostatic term, the complicated computations involved in model II are avoided. In the limiting case, if  $n_{\rm B}=0$  in model III, the free-energy change given by equation (14) is the same as that given by equation (6), which has been shown to agree well with other more rigorous calculations, especially those of Briant and Burton (ref. 11). Computed cluster spectra for the three models under various relative-acidity and relative-humidity combinations are given in tables 1 through 3.

Figure 3 shows the spectra of calculated ion- $H_2O-H_2SO_4$  clusters for the three models (Relative humidity = 0.1 and Relative acidity = 1 × 10<sup>-5</sup>). The value of the  $H^{-4}O$  ion radius in these calculations is assumed to be 1.38 Å. For model I, the ion clusters are mainly confined to groups containing one or two sulfuric acid molecules with a relatively small number of water molecules attached to the ion core. For model II, the ion-cluster spectrum is spread out, and ion clusters containing higher numbers of sulfuric acid and water molecules are more common. For model III, there is an appreciable amount of the group containing three sulfuric acid molecules accompanied by further spectral broadening. The peak intensity (relative abundance) of the cluster distribution is lowered while the spectrum gets broader. Table 4 shows that the mean radius of the ion clusters increases from model I to model II to model III. Hence, the effect of the modifications included in model II over the conventional extended Thomson theory (model I) is to broaden the ion-cluster spectrum, producing a more even distribution and bigger ion clusters. This results in a larger mean radius in accord with the experimental results reported by Singh et al. (See ref. 23.) These features are further strengthened in model III.

<sup>\*</sup>For dilute clusters, the g factors are approximately the same as for water. 
\*\*The optimum value of  $\gamma$  for ion-H<sub>2</sub>O clusters is 1.3. The same value of  $\gamma$  has been assumed for ion-H<sub>2</sub>O-H<sub>2</sub>SO<sub>4</sub> clusters.

#### CONCLUDING REMARKS

Examination of the three models reveals that the microscopic surface tension used in models II and III greatly reduces the surface-energy contribution for small ion clusters and hence enhances the formation of ultrafine ion clusters. Comparison of models I, II, and III also shows that the electrostatic-energy term is important. A slight increase in the effective charge of the ion facilitates the formation of larger and more stable prenucleation clusters. The effect of the neutral sulfuric acid hydrates is small, especially when the relative humidity is low.

From the comparison of various models for various combinations of relative humidity and relative acidity, it can be seen that the Thomson theory (model I) consistently predicts smaller clusters with narrowly peaked distributions, whereas the modified theories (models II and III) predict larger clusters with broader distributions. These latter trends are consistent with recently reported experimental results.

The modifications involving the curvature dependence of the microscopic surface tension and strong intermolecular interaction between the ion core and its surrounding molecules introduce large differences between the predicted spectral distributions of the ion-H<sub>2</sub>O-H<sub>2</sub>SO<sub>4</sub> clusters for the various models. Because of the higher contribution of the electrostatic-energy term in the present model (model III), the predicted microcluster spectrum becomes even broader, and more of the larger ion clusters are included. The validity of the charge scaling-factor model has yet to be tested. However, the agreement in the values of free energy of formation of F and Cs hydrates in the present model with other, more rigorous, theories suggests that it may be a reasonably accurate model to use for free-energy calculations in the region of very small clusters.

Langley Research Center National Aeronautics and Space Administration Hampton, VA 23665 June 15, 1982

#### REFERENCES

- 1. Arnold, F.; Henschen, G.; and Ferguson, E. E.: Mass Spectrometric Measurements of Fractional Ion Abundances in the Stratosphere Positive Ions. Planet. & Space Sci., vol. 29, 1981, pp. 185-193.
- Arnold, F.; Fabian, R.; Ferguson, E. E.; and Joos, W.: Mass Spectrometric Measurements of Fractional Ion Abundances in the Stratosphere Negative Ions.
  Planet & Space Sci., vol. 29, 1981, pp. 195-203.
- 3. Singh, Jag J.; and Khandelwal, G. S.: Elemental Characteristics of Aerosols Emitted From a Coal-Fired Heating Plant. NASA TM-78749, 1978.
- 4. Thomson, J. J.; and Thomson, G. P: Conduction of Electricity Through Gases.

  Volume I General Properties of Ions Ionisation by Heat and Light, Third ed.

  Cambridge Univ. Press, 1928.
- 5. Castleman, A. W., Jr.: Studies of Ion Clusters: Relationship to Understanding Nucleation and Solvation Phenomena. Kinetics of Ion-Molecule Reactions, Pierre Ausloos, ed., Plenum Press, Inc., c.1979, pp. 295-321.
- 6. Castleman, A. W., Jr.: Nucleation and Molecular Clustering About Ions. Adv. Colloid & Interface Sci., vol. 10, 1979, pp. 73-128.
- 7. Castleman, A. W., Jr.: A Reconsideration of Nucleation Phenomena in Light of Recent Findings Concerning the Properties of Small Clusters, and a Brief Review of Some Other Particle Growth Processes. Astrophys. & Space Sci., vol. 65, no. 2, Oct. 1979, pp. 337-349.
- 8. Castleman, A. W.: Neutral and Charged Clusters in the Atmosphere: Their Importance and Potential Role in Heterogeneous Catalysis. Heterogeneous Atmospheric Chemistry, David R. Schryer, ed., Geophys. Monogr. Ser., vol. 26, American Geophys. Union, 1982.
- 9. Chan, Lo Yin; and Mohnen, V. A.: Ion Nucleation Theory. J. Atmos. Sci., vol. 37, no. 10, Oct. 1980, pp. 2323-2331.
- 10. Burton, J. J.: Nucleation Theory. Statistical Mechanics Part A: Equilibrium Techniques, Bruce J. Berne, ed., Plenum Press, c.1977, pp. 195-234.
- 11. Briant, C. L.; and Burton, J. J.: A Molecular Model for the Nucleation of Water on Ions. J. Atmos. Sci., vol. 33, no. 7, July 1976, pp. 1357-1361.
- 12. Singh, Jag J.; Smith, Alphonso C.; Chan, Lo Y.; and Yue, Glenn K.: Calculation of Composition Distribution of Ultrafine Ion-H<sub>2</sub>O-H<sub>2</sub>SO<sub>4</sub> Clusters Using a Modified Binary Ion-Nucleation Theory. NASA TP-2031, 1982.
- 13. Tolman, Richard C.: The Effect of Droplet Size on Surface Tension. J. Chem. Phys., vol. 17, no. 3, Mar. 1949, pp. 333-337.
- 14. Kim, Myungkun; and Chang, Seihun: Effect of the Curvature Dependency of Surface Tension of Pendular Ring Condensation Calculations. J. Colloid & Interface Sci., vol. 69, no. 1, 1979, pp. 34-44.

- 15. Vogelsberger, Wolfram: Influence of Curvature-Dependent Surface Tension on the Free Energy of Formation of Microclusters. Chem. Phys. Lett., vol. 74, no. 1, Aug. 15, 1980, pp. 143-146.
- 16. Suck, Sung Ho: Change of Free Energy in Heteromolecular Nucleation: Electrostatic Energy Contribution. J. Chem. Phys., vol. 75, no. 10, Nov. 15, 1981, pp. 5090-5096.
- 17. Yue, Glenn K.; and Chan, L. Y.: Theory of the Formation of Aerosols of Volatile Binary Solutions Through the Ion-Induced Nucleation Process. J. Colloid & Interface Sci., vol. 68, no. 3, Mar. 1979, pp. 501-507.
- 18. Suzuki, Kunihiro; and Mohnen, Volker A.: Binary Homogeneous Nucleation Theory by Cluster-Cluster Interaction With Application to the H<sub>2</sub>SO<sub>4</sub>-H<sub>2</sub>O System. J. Aerosol Sci., vol. 12, no. 1, 1981, pp. 61-73.
- 19. Liszi, J.; Mészáros, L.; and Ruff, I.: The Field Dependence of the Kirkwood Factor and the Nonlinear Dielectric Behavior of Some Liquids. J. Chem. Phys., vol. 74, no. 12, June 15, 1981, pp. 6896-6901.
- 20. Kirkwood, John G.: The Dielectric Polarization of Polar Liquids. J. Chem. Phys., vol. 7, no. 10, Oct. 1939, pp. 911-919.
- 21. Sabinina, L.; and Terpugov, L.: The Surface Tension of the System Sulfuric Acid-Water. Z. Phys. Chem. (Neue Folge), vol. A173, 1935, pp. 237-241.
- 22. Ayers, G. P.; Gillett, R. W.; and Gras, J. L.: On the Vapor Pressure of Sulfuric Acid. Geophys. Res. Lett., vol. 7, no. 6, June 1980, pp. 433-436.
- 23. Singh, J. J.; Smith, A. C.; and Yue, G. K.: Experimental Studies of the Ion-Induced Binary Nucleation. J. Aerosol Sci., vol. 13, no. 4, July 1982.

TABLE 1.- COMPARISON OF PREDICTED ION-CLUSTER DISTRIBUTIONS FOR VARIOUS MODELS

WITH R.A. =  $1 \times 10^{-6}$  AND R.H. = 0.1

Only ion-cluster intensities greater than 1 percent in the spectrum are listed

Model			P(	n <sub>B</sub> ,n <sub>A</sub> )	for n	<sub>B</sub> ,n <sub>A</sub> v	alues o	f -			$\sum_{n_{A}} P(n_{B}, n_{A})$
	1,2	1,3	1,4	1,5	1,6	1,7	1,8	1,9	1,10	1,11	(1,x)
III		2.5%	19.7% 5.0%	33.7% 17.1%	22.3% 20.7% 2.9%	8.1% 12.6% 4.3%	1.8% 4.5% 3.3%	1.1% 1.6%			88.1% 61.0% 12.1%
	2,2	2,3	2,4	2,5	2,6	2,7	2,8	2,9	2,10	2,11	(2,x)
III				2.9% 4.4% 2.3%	4.1% 11.2% 11.6%	2.4% 11.3% 21.7%	6.9% 22.8%	2.8% 14.9%	6.0%	2.1%	9.4% 36.6% 81.4%

TABLE 2.- COMPARISON OF PREDICTED ION-CLUSTER DISTRIBUTIONS FOR VARIOUS MODELS

WITH R.A. =  $1 \times 10^{-4}$  AND R.H. = 0.1

Only ion-cluster intensities greater than 1 percent in the spectrum are listed

Model					P(n <sub>B</sub> ,	n <sub>A</sub> ) fo	r n <sub>B</sub> ,n	A valu	es of -					$\sum_{n_{A}} P(n_{B}, n_{A})$
	1,2	1,3	1,4	1,5	1,6	1,7	1,8	1,9	1,10	1,11	1,12	1,13	1,14	(1,x)
I II III			1.4%	2.3%	1.6%									5•3%
	2,2	2,3	2,4	2,5	2,6	2,7	2,8	2,9	2,10	2,11	2,12	2,13	2,14	(2,x)
I II III			6•6%.	20.2% 6.0%	28.2% 15.3% 1.7%	16.6% 15.6% 3.2%	6.1% 9.5% 3.4%	1.5% 3.8% 2.2%						79.2% 50.2% 10.5%
	3,2	3,3	3,4	3,5	3,6	3,7	3,8	3,9	3,10	3,11	3,12	3,13	3,14	(3,x)
I II III					2.6% 3.2% 1.5%	3.7% 7.7% 5.6%	3.1% 10.9% 11.8%	1.9% 10.7% 16.8%	5.2% 11.8%	2.2% 6.9%	2.4%			11.3% 39.9% 56.8%
	4,2	4,3	4,4	4,5	4,6	4,7	4,8	4,9	4,10	4,11	4,12	4,13	4,14	(4,x)
I II III							1.5%	3.5%	5.5%	6.6%	4.8%	2.3%	1.1%	25•3%

#### TABLE 3.- COMPARISON OF PREDICTED ION-CLUSTER DISTRIBUTIONS FOR VARIOUS MODELS

WITH R.A. =  $1 \times 10^{-6}$  AND R.H. = 0.5

Only ion-cluster intensities greater than 1 percent in the spectrum are listed

Model				-		P	(n <sub>B</sub> ,n <sub>A</sub> )	for	n <sub>B</sub> ,n <sub>A</sub>	values	of -							$\sum_{n_{A}} P(n_{B}, n_{A})$
	1,4	1,5	1,6	1,7	1,8	1,9	1,10	1,11	1,12	1,13	1,14	1,15	1,16	1,17	1,18	1,19	1,20	(1,x)
III		2.1%	6.9%	12.6%	13.9% 2.8%	10.8% 3.5%	6.9%	3.5% 2.5%	1.6%									58.3% 15.3%
	2,4	2,5	2,6	2,7	2,8	2,9	2,10	2,11	2,12	2,13	2,14	2,15	2,16	2,17	2,18	2,19	2,20	(2,x)
III II	·		1.3%	3.7% 1.4%	6.8% 4.4% 2.2%	8.5% 8.8% 4.4%	7.1% 11.1% 7.6%	13.1%	3.3% 10.9% 10.5%	1.8% 8.6% 8.6%	5.1% 6.7%	2.9% 4.5%	1.5% 2.6%	1.4%				38.1% 67.8% 57.9%
	3,4	3,5	3,6	3,7	3,8	3,9	3,10	3,11	3,12	3,13	3,14	3,15	3,16	3,17	3,18	3,19	3,20	(3,x)
III								1.3% 1.2%	1.7%	2.2% 3.7%	2.0% 4.4%	1.7%		3.8%	2.8%	2.0%	1.1%	10.2% 30.7%

TABLE 4.- MEAN RADII OF ION-CLUSTER SPECTRA FOR VARIOUS MODELS

WITH R.H. = 0.1

Mean radius = 
$$\bar{r} = \frac{\sum_{i}^{p_i r_i}}{\sum_{i}^{p_i}}$$
 when  $P_i$  is the probability for radius  $r_i$ 

	м	ean radius	for R.A.	values of	_
Model	1 × 10 <sup>-6</sup>	1 × 10 <sup>-5</sup>	1 × 10 <sup>-4</sup>	1 × 10 <sup>-3</sup>	1 × 10 <sup>-2</sup>
III	3.67 Å 3.92 Å 4.29 Å	3.90 Å 4.18 Å 4.47 Å	4.15 Å 4.45 Å 4.83 Å	4.51 Å 5.04 Å 5.50 Å	5.77 Å 7.25 Å 7.35 Å

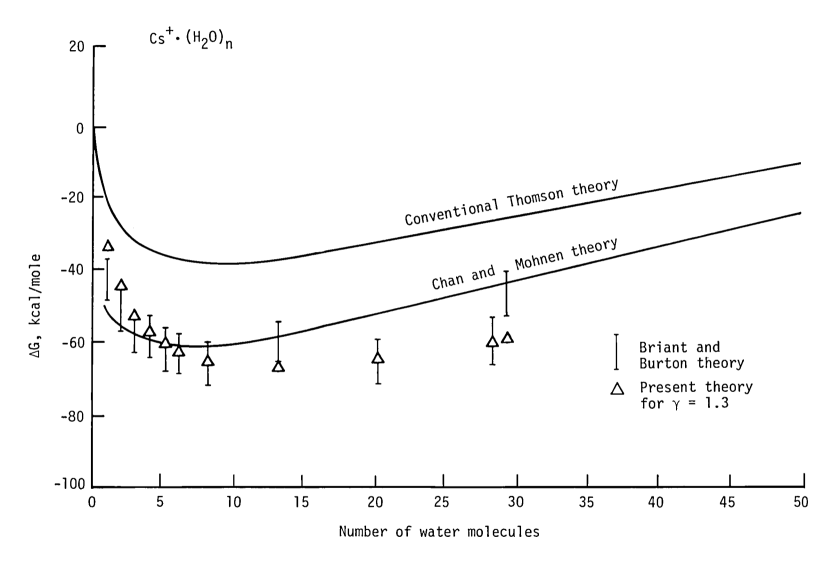


Figure 1.- Comparison of  $\Delta G$  for four different models at S=1 for  $Cs^+ \cdot (H_2O)_n$  clusters.

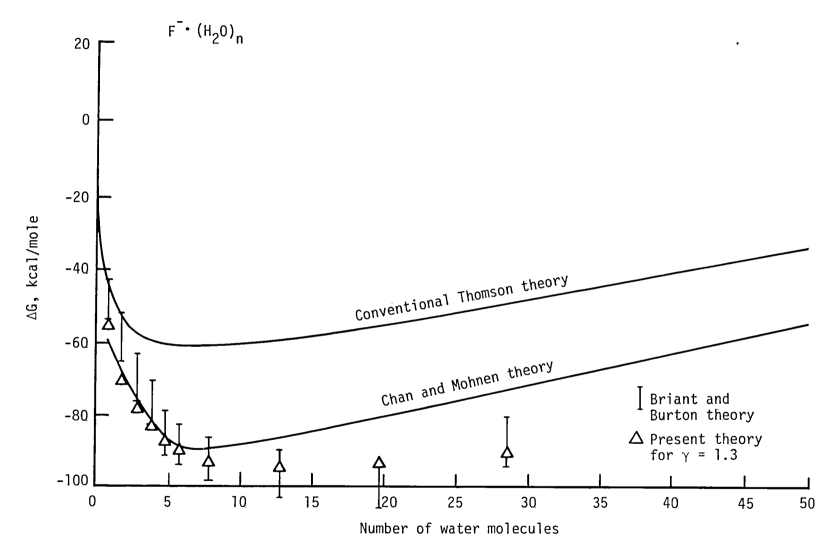


Figure 2.- Comparison of  $\Delta G$  for four different models at S=1 for F  $\bullet$  (H2O) $_n$  clusters.

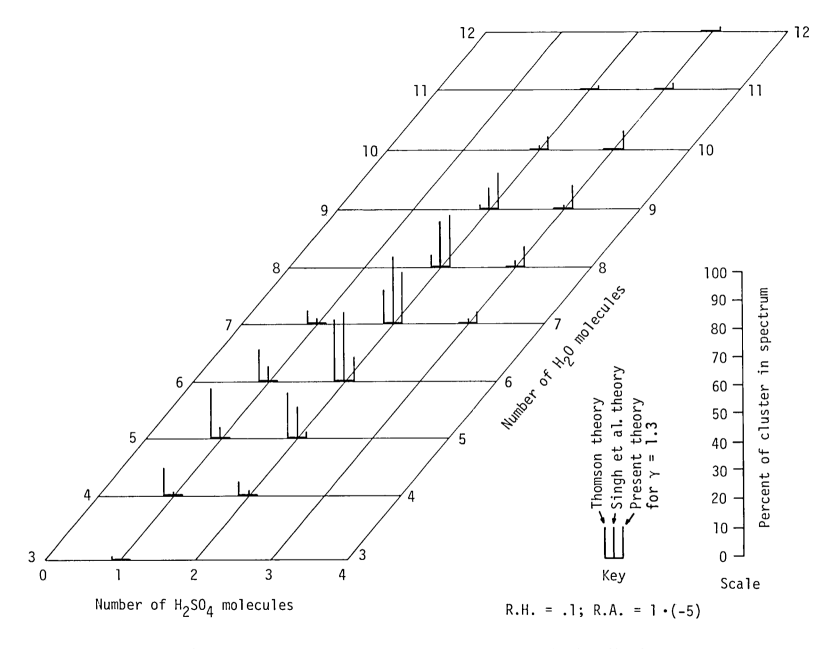


Figure 3.- Comparison of ion- ${\rm H_2O-H_2SO_4}$  spectral distribution for three different models.

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